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Clinical Pharmacology 1: Phase 1 Studies and Early Drug Development

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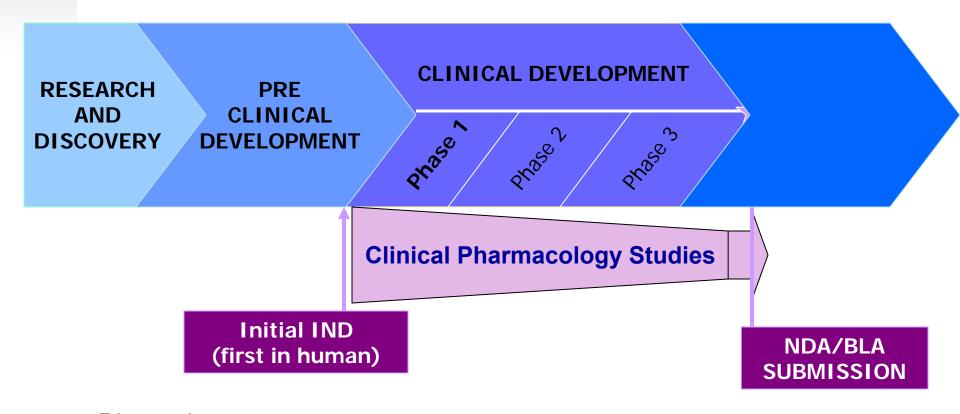


Objectives

- Outline the Phase 1 studies conducted to characterize the Clinical Pharmacology of a drug; describe important design elements of and the information gained from these studies.
- List the Clinical Pharmacology characteristics of an Ideal Drug
- Describe how the Clinical Pharmacology information from Phase 1 can help design Phase 2/3 trials
- Discuss the timing of Clinical Pharmacology studies during drug development, and provide examples of how the information generated could impact the overall clinical development plan and product labeling.



Phase 1 of Drug Development



Phase 1

 studies designed mainly to investigate the safety/tolerability (if possible, identify MTD), pharmacokinetics and pharmacodynamics of an investigational drug in humans

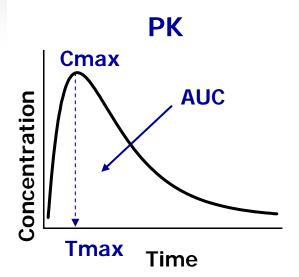


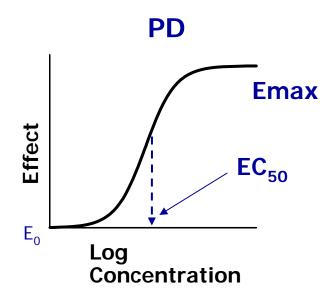


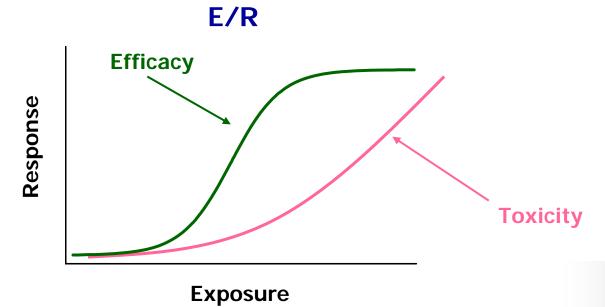
Clinical Pharmacology

- Study of the Pharmacokinetics (PK) and Pharmacodynamics (PD) of the drug in humans
 - PK: what the body does to the drug (<u>Absorption</u>,
 <u>Distribution</u>, <u>Metabolism</u>, <u>Excretion</u>)
 - PD: what the drug does to the body
- PK and PD profiles of the drug are influenced by physicochemical properties of the drug, product/formulation, administration route, patient's intrinsic and extrinsic factors (e.g., organ dysfunction, diseases, concomitant medications, food)





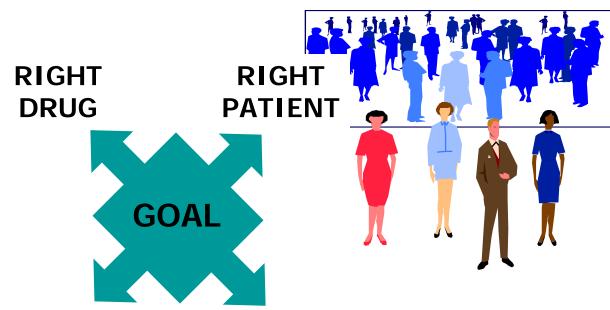






The Ultimate Goal:





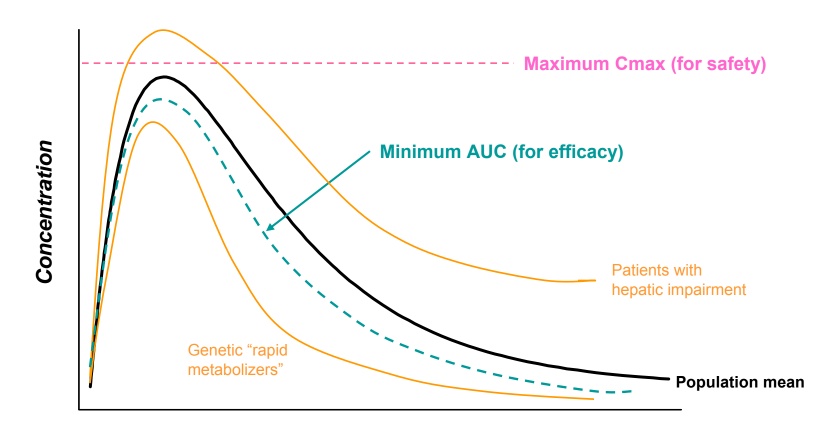
RIGHT DOSE







To determine the dose/dosing regimen that achieves target drug exposures <u>in all</u> <u>relevant populations</u>







How do we achieve the goal?

Clinical Pharmacology

- First-in-Human
- SAD and MAD PK Studies
- Healthy vs. Patient population
- ADME (Mass Balance)
- Specific Populations
 - Renal Impairment
 - Hepatic Impairment
 - Age, gender, etc.
 - Pediatrics
- Drug Interactions
- Population PK
- Biomarkers
- Pharmacogenomics
- Special Safety (e.g., TQTc study)

Exposure-response (PK/PD)

- Dose selection and optimization
- Efficacy vs. Safety
- Quantitative approaches
 - Clinical trial simulation
 - Disease models

Biopharmaceutics

- Bioavailability/Bioequivalence (BA/BE)
- Food Effect

In Vitro Studies

- Protein Binding
- Blood to Plasma Partitioning
- In vitro drug metabolism, transport and drug interactions

Bioanalytical Methods

Assay Validation & Performance Reports

Biologics only

- Immunogenicity
- Comparability





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Single Dose/Multiple Dose Escalation Studies

- Typically the first-in-human study (or studies)
- Randomized, placebo-controlled, healthy volunteers (or patients, in certain cases)
- Starting dose determined by preclinical toxicology studies
- Information gained:
 - Safety/tolerability, identify maximum tolerated dose (MTD)
 - General PK characteristics, variability, linearity/ proportionality
 - Steady-state parameters (accumulation, time-dependency)
 - Preliminary exploration of drug elimination (urine PK, metabolite identification)





ADME (i.e. Mass Balance) Study*

- Objective: To understand the full clearance mechanisms of the drug and its metabolites in humans
- Typically single dose, healthy males (n=4-6), at intended route of administration
- Radio-labeled (C¹⁴) drug molecule
- Measure concentrations of parent and metabolite(s) and determine amt of radioactivity in plasma, urine, feces
- Information gained:
 - Primary mechanism(s) of elimination and excretion from the body
 - Proportion of parent drug converted to metabolite(s)



^{*} Not usually done with high MW therapeutic proteins

BA/BE Studies

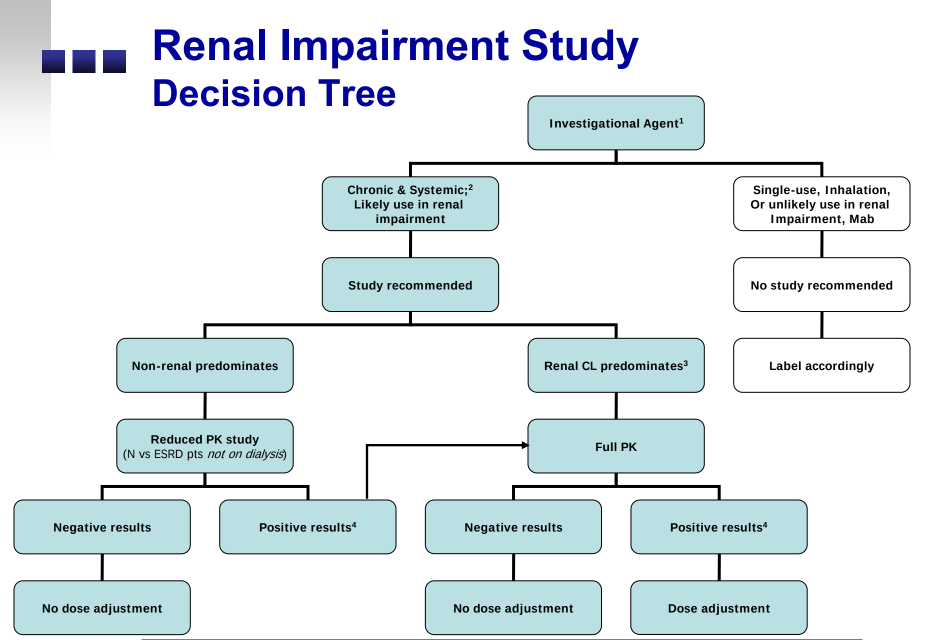
- Objective: To evaluate the rate (Cmax, Tmax) and extent (AUC) of absorption of drug from a test formulation (vs. reference formulation)
- BA:Typically, crossover, single dose (if linear PK) study in healthy subjects; measure blood/plasma conc. of parent drug and major active metabolites for ≥ 3 t½
 - **BE**: crossover study in fasted healthy subjects given <u>single</u> <u>doses</u> of test & reference products administered <u>at same molar doses</u>; measure blood/plasma conc. of parent drug only
- "Pivotal" BE study required to bridge the to-be-marketed formulation (test) to that used in Phase 3 clinical trials (reference)
- BE acceptance criteria: 90% CI of test/reference Cmax & AUC ratios within 80-125%
- Information gained:
 - Relative BA, Absolute BA of drug from a formulation
 - BE (no significant difference in BA) of test vs. reference



Food Effect Study

- Objective: To evaluate the effect of food on rate and extent of drug absorption from a given formulation
- Single dose, crossover, two-treatment (fed vs fasted), two-period, two-sequence study in healthy subjects (n ≥ 12 with data); use highest strength of drug product; fed: FDA high-fat high-calorie meal
- PK assessments similar to BA study
- No food-effect if 90% CI of fed/fasted Cmax and AUC ratios within 80-125%. The clinical significance of any observed food effect could be determined based on drug's exposure-response profile.
- Information gained:
 - effect of food on the BA of oral drugs
 - Labeling instructions on whether to administer drug on empty stomach or without regard to meals







¹Metabolites (active/toxic) – same decision tree

4 >50% increase in AUC; < for Narrow TI drugs

² Includes cytokines or cytokine modulators with MW <69 kDa

³ Option to do either full or reduced study or Pop PK Analysis of Ph 2/3 data

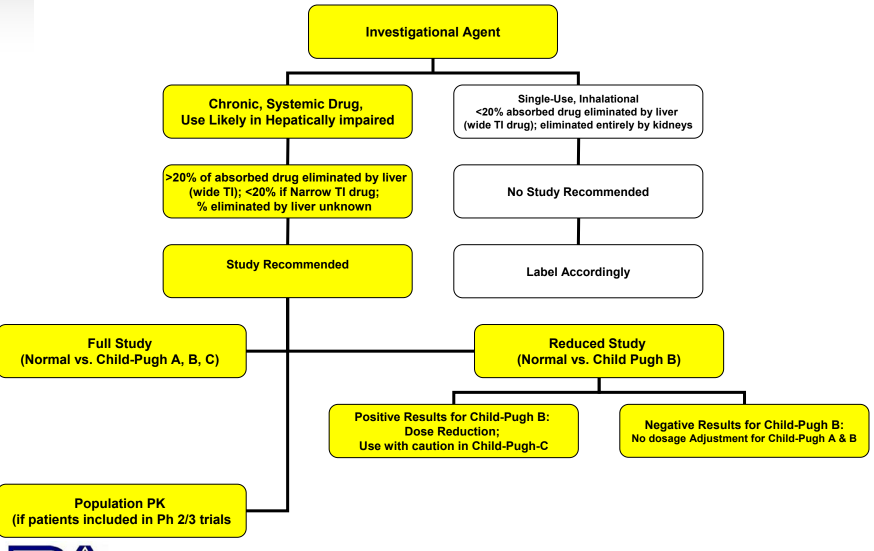
Renal Impairment Study Full Study Design

- Single dose (if linear & time independent PK), parallel groups, "healthy" males and females with varying degrees of renal function (≥6 per group)
- Calculate CrCl via Cockcroft-Gault; eGFR via MDRD
- Stratification (based on CrCl): Normal (≥90 mL/min), Mild (60-89 mL/min), Moderate (30-59 mL/min) and Severe Impairment (15-29 mL/min), ESRD (<15 mL/min) dialysis and non-dialysis
- Information gained:
 - Effect of renal impairment on drug clearance; dosage recommendations for various stages of renal impairment
 - Effect of hemodialysis (HD) on drug exposure; info on whether dialysis could be used as treatment for drug overdosage





Hepatic Impairment Study Decision Tree





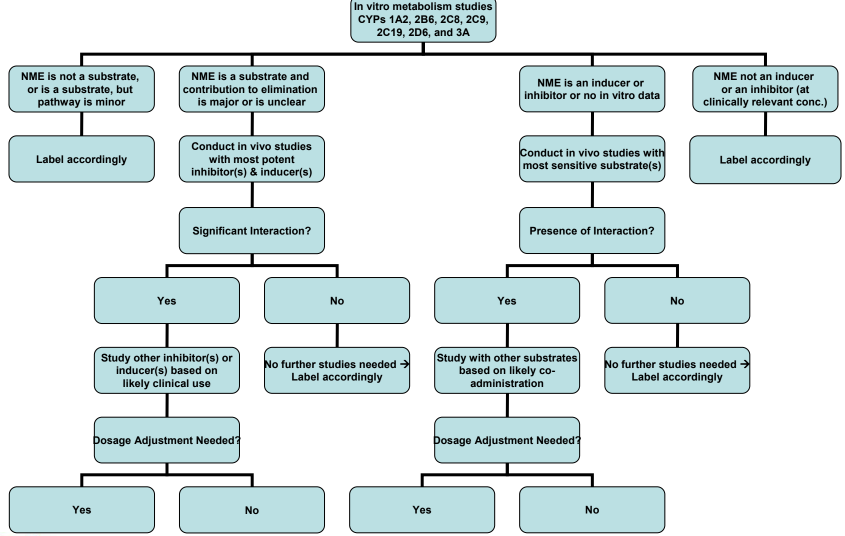


Hepatic Impairment Study

- Study Designs:
 - (1) Full Study: Single dose (if linear & time-independent PK), parallel groups, males & females with varying degrees of hepatic impairment (≥6 per group)
 - Normal Hepatic Function (matched for age, gender & BW to subjects with hepatic impairment)
 - Child-Pugh Class A (Mild)
 - Child-Pugh Class B (Moderate)
 - Child-Pugh Class C (Severe)
 - (2) Reduced Study: Normal vs. Child-Pugh B (Moderate) (≥8 per group)
 - (3) Pop-PK approach
- If drug is metabolized by enzyme with genetic polymorphisms (e.g. CYP2C19, CYP2D6), genotype status of subjects should be assessed and considered during PK data analysis.
- Information gained:
 - Effect of hepatic impairment on PK of parent drug and metabolites
 - Dosage recommendations for various stages of hepatic impairment



Drug Interaction Studies Decision Tree for CYP450





Drug Interaction Studies

- Objective: To evaluate potential of investigational drug as an inhibitor/inducer (I) and substrate (S) of certain metabolizing enzymes/transporters
- Preferably crossover design (parallel if long t½ drug); healthy subjects (or patients for safety considerations or if desirable to evaluate PD endpoints)
- The choice of doses/dosing intervals/dosage forms of substrate and inhibitor/inducer, routes & timing of coadministration, number of doses should maximize possibility of finding an interaction and mimic the clinical setting, with due consideration for safety of study population.
- Degree of effect (inhibition/induction) is typically classified by change in the substrate AUC:
 - e.g., Drug causes ≥ 5-fold increase in midazolam AUC → "potent" inhibitor of CYP3A4
- Exposure-response information on the drug is important in assessing the clinical significance of the change in AUC of substrate by inhibitor/inducer.





Thorough QT Study (TQT)

- In vivo safety study required for all systemically available NMEs (regardless of in vitro or non-clinical findings)
- Objective: To identify drugs that prolong QT(95% CI upper bound ≥ 10 ms) that need a more thorough ECG monitoring in pivotal trials; TQT study conducted prior to Phase 3 trials
- Usually, single dose study in healthy subjects; evaluate therapeutic and "supratherapeutic" doses of drug versus positive control (e.g., moxifloxacin)
- ICH Guidelines, E14: The Clinical Evaluation of QT/QTc Interval Prolongation and Proarrhythmic Potential for Non-Antiarrhythmic Drugs
 - Recommendations for design, conduct, analysis, and interpretation of clinical studies



ABSORPTION:

- High absolute bioavailability with low variability
- Exhibits linear PK over therapeutic dose range, i.e. dose-proportional increases in Cmax, AUC
 - Single-dose study design sufficient: BA, PK in renal impairment, hepatic impairment & DDI
- AUC, Cmax not significantly affected by concomitant food, pH-altering medications, grapefruit, alcohol, etc.
- BCS Class I (high solubility + high permeability)
 - can qualify for biowaiver of future additional BA/BE studies





DISTRIBUTION:

- Reaches the target site(s) of action immediately and at effective/nontoxic concentrations; doesn't accumulate in non-target organs
 - Local (targeted) application advantageous over systemic administration
- Not significantly (>80 to >95%) bound to plasma proteins; extent of PB not concentration- and timedependent
 - only free or unbound drug is active
 - less prone to DDI with highly-protein drugs (e.g., warfarin)
 - may require that plasma samples be assayed for the free drug (e.g., in PK studies in renal and hepatic impairment)





METABOLISM/EXCRETION:

- Not extensively metabolized or not exclusively metabolized by a CYP450 enzyme
 - CL less likely to be affected by hepatic impairment and/or concomitant administration of other drugs that affect one or more metabolizing enzymes
- Not metabolized by polymorphic enzymes (e.g., CYPs 2D6, 2C19, 2C9, NAT2)
 - does not require genotyping in PK and other clinical studies
- CL not highly variable depending on 'covariates' as age, race, gender, disease/comorbidities
- CL not time-dependent (e.g., metabolic autoinduction, diurnal variation)
 - may require longer duration of studies for PK profiling



OTHERS:

- Not a Narrow Therapeutic Index Drug
 - slight changes in drug exposure less likely to impact efficacy/safety
 - less likely to require therapeutic drug monitoring in clinical trials and clinical practice to minimize toxicities and lack of efficacy
- Does not prolong the QT interval
 - less likely to have TdP risk
- Not a significant inhibitor or inducer of CYP3A, P-gp, etc.
 - less likely to have DDI with concomitantly administered drugs
- Does not trigger formation of neutralizing anti-drug antibodies or organ-damaging immune complexes (immunogenicity)

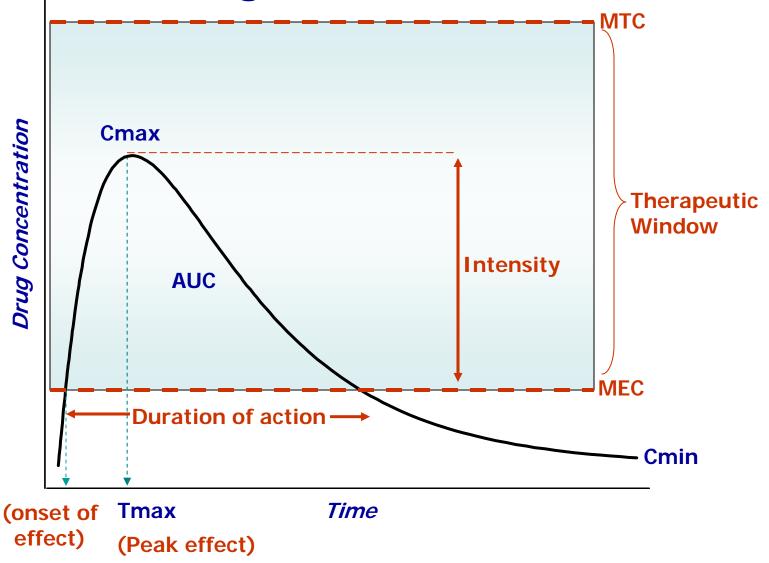
PK Parameters and Design of Phase 2/3 Trials

Parent Drug and Active Metabolites:

- T_{max}
 -represent the most appropriate time(s) to perform safety assessments (e.g., vital signs, ECG, other immediate PD effects)
- $t^{1/2}$
 - considered when determining dosage interval
 - related to time to steady state (t_{ss}) after dose initiation or dose adjustment; considered in evaluating need for a loading dose
 - influences the duration of monitoring after dosing and follow-up after withdrawal of therapy
 - determines adequate washout period between treatments (in crossover studies)
- Cmax, Cmin, AUC
 - important for dose selection (viewed relative to MEC and MTC)
 eg. PK/PD parameters predicting efficacy of anti-infectives

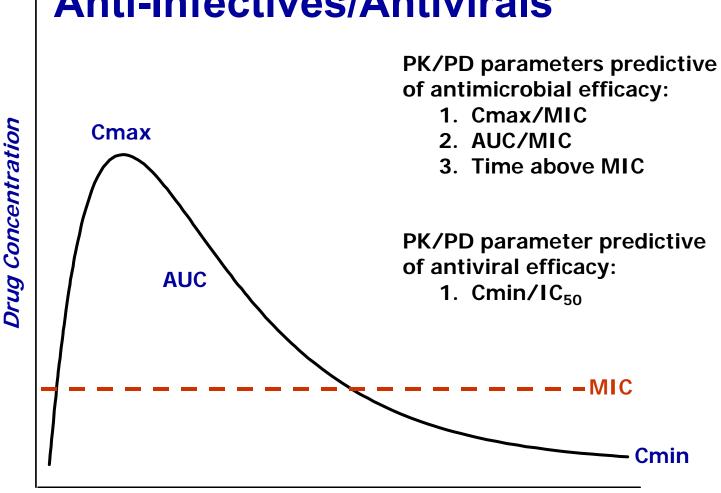


PK and Drug Effect





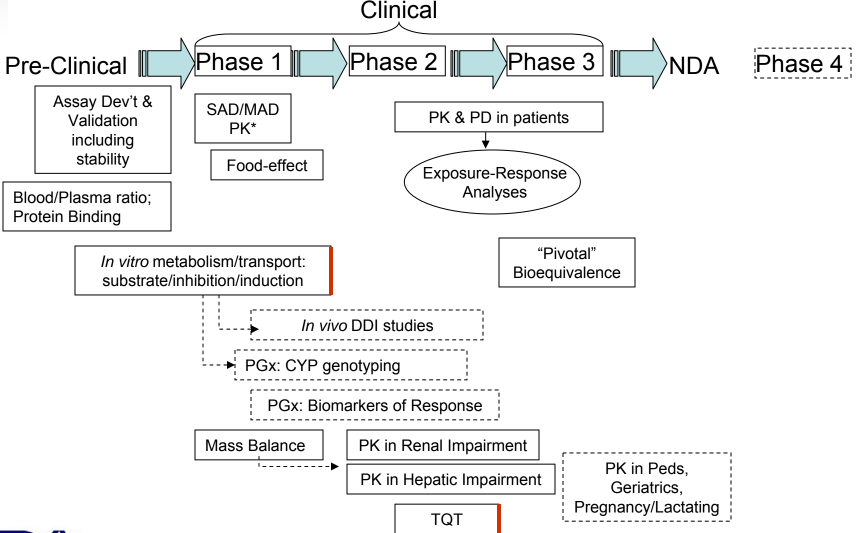
PK and Drug Effect EXAMPLE: Anti-Infectives/Antivirals







Timing of Early and Clinical Pharmacology Studies







Phase 1 Studies: Impact on Labeling

FULL PRESCRIBING INFORMATION:

- 1 INDICATIONS AND USAGE
- 2 DOSAGE AND ADMINISTRATION
- **3 DOSAGE FORMS AND STRENGTHS**
- **4 CONTRAINDICATIONS**
- **5 WARNINGS AND PRECAUTIONS**
- **6 ADVERSE REACTIONS**
- 7 DRUG INTERACTIONS
- **8 USE IN SPECIFIC POPULATIONS**
 - 8.1 Pregnancy
 - 8.2 Labor and Delivery
 - 8.3 Nursing Mothers
 - 8.4 Pediatric Use
 - 8.5 Geriatric Use

9 DRUG ABUSE AND DEPENDENCE

- 9.1 Controlled Substance
- 9.2 Abuse
- 9.3 Dependence
- **10 OVERDOSAGE**
- 11 DESCRIPTION

12 CLINICAL PHARMACOLOGY

- 12.1 Mechanism of Action
- 12.2 Pharmacodynamics
- 12.3 Pharmacokinetics
- 13 NONCLINICAL TOXICOLOGY
- 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility
- 13.2 Animal Toxicology and/or Pharmacology
- 14 CLINICAL STUDIES
- 15 REFERENCES
- 16 HOW SUPPLIED/STORAGE AND HANDLING
- 17 PATIENT COUNSELING INFORMATION





Clinical Pharmacology Guidance Documents

- Clinical Lactation Studies (2005*)
- Clinical Pharmacogenomics (2011*)
- Drug Interaction Studies (2006*,1999,1997)
- Drug Metabolism/Drug Interaction Studies in the Drug Development Process: Studies In Vitro (1997)
- General Considerations for Pediatric Pharmacokinetic Studies for Drugs and Biological Products (1998*)
- In Vivo Drug Metabolism/Drug Interaction Studies (1999)
- Pharmacokinetics in Patients with Impaired Hepatic Function (2003)
- Pharmacokinetics in Patients with Impaired Renal Function (2010*, 1998)
- Pharmacokinetics in Pregnancy (2004*)
- Population Pharmacokinetics (1999)
- Exposure-Response Relationships Study Design, Data Analysis, and Regulatory Applications (2003)





- Bioanalytical Method Validation (2001)
- Bioavailability and Bioequivalence Studies for Nasal Aerosols and Nasal Sprays for Local Action (2003*)
- Bioavailability and Bioequivalence Studies for Orally Administered Drug Products (2003)
- Dissolution Testing of Immediate Release Solid Oral Dosage Forms (1997)
- Extended Release Oral Dosage Forms: Development, Evaluation, and Application of In Vitro/In Vivo Correlations (1997)
- Food-Effect Bioavailability and Fed Bioequivalence Studies (2002)
- Statistical Approaches to Establishing Bioequivalence (2001)
- Waiver of In Vivo Bioavailability and Bioequivalence Studies for Immediate-Release Solid Oral Dosage Forms Based on a Biopharmaceutics Classification System (2000)

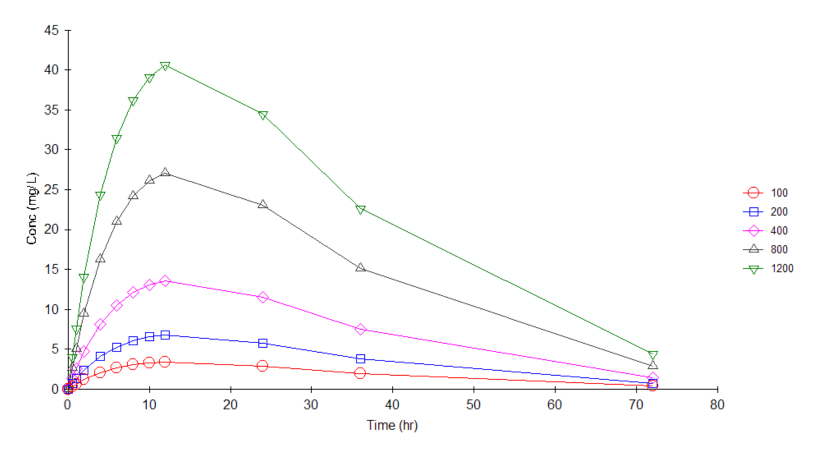


EXAMPLES





■■■ SAD PK Study Results (Example)





Food Effect Example: REYATAZ® (atazanavir) oral capsules

- Administration of a single dose of atazanavir (800 mg) with a light meal increased Cmax by 57% and AUC by 70%; a high-fat meal increased AUC by 35% with no change in Cmax. The % CVs of AUC and Cmax decreased by approximately one-half compared to the fasting state.
- Clinical trials were conducted under fed conditions.
- Label directs administration with a meal or snack.



■■■ Renal Impairment Example: DORIBAX® (doripenem) powder for IV use

- In a radiolabeled ADME study, approximately 93% of the dose was excreted in the urine by 12 hours. Less than 1% of the total radioactivity was recovered in feces after one week.
- Because doripenem is primarily eliminated by the kidneys, a Full PK study in patients with renal impairment was conducted.
- In Phase 2/3 trials, dosage was adjusted based on CrCL.
- The label recommends dosage reduction for patients with moderate or severe renal impairment... and hemodialysis as a treatment for overdosage.





- In vitro metabolism studies using human liver microsomes indicated that raltegravir is not a substrate of CYP450 enzymes but is metabolized mainly by UGT1A1. A Mass Balance study showed that Raltegravir is eliminated primarily by glucuronidation in the liver. Renal clearance is a minor pathway of elimination.
- In the PK-Hepatic Impairment Study (Reduced Study Design), there were no clinically important pharmacokinetic differences between subjects with moderate hepatic impairment and healthy subjects.
- PopPK analysis of Phase 2/3 trial data further indicates that the PK of raltegravir in Child Pugh B were not different from patients with normal hepatic function.
- Labeling states: No dosage reduction for patients with moderate or <u>mild hepatic</u> impairment is recommended. The effect of severe <u>hepatic</u> impairment on the PK of the drug was not studied.



